

# TeraChem User's Guide

Version 1.5K

PetaChem, LLC  
26040 Elena Road  
Los Altos Hills, CA 94022

<http://www.petachem.com>

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## I Introduction

TeraChem is general purpose quantum chemistry software designed to run on Nvidia CUDA-enabled GPU architectures under a 64-bit Linux operating system. Some of TeraChem features include:

- Restricted, unrestricted, and restricted open shell Hartree-Fock and grid-based Kohn-Sham energy and gradient calculations
- Full support of s, p and d-type atom-centered Gaussian basis functions
- Various DFT functionals, including range-corrected and Coulomb attenuated functionals (BLYP, B3LYP, PBE, PBE0,  $\omega$ PBE,  $\omega$ PBEh,  $\omega$ B97,  $\omega$ B97x, camB3LYP, etc) and DFT grids (800 - 80,000 grid points per atom)
  - Static grid (single grid used for the entire calculation) and dynamical grid (multigrid) integration.
  - Empirical dispersion correction (DFT-D3 and DFT-D2)
- Geometry optimization (L-BFGS, Conjugate gradient, Steepest descent)
  - The optimization can be carried out either in Cartesian or internal coordinates as specified in the start file (all input geometries are provided in Cartesians). The Cartesian → internal → Cartesian coordinate transformation is performed automatically whenever required.
  - Constrained optimization with frozen atoms, constrained bond lengths, angles, and dihedrals.
- Transition state search (Nudged elastic band) in internal and Cartesian coordinates
- Ab initio molecular dynamics (NVE, NVT ensembles)
  - Reversible Born-Oppenheimer dynamics
  - Spherical boundary conditions
- Support of multiple-GPU systems
- Single/Dynamical/Double precision accuracy
- QM/MM treatment of surrounding water molecules using TIP3P force field<sup>1</sup>
- Natural bond orbital analysis through integration with NBO6
- Polarizabilities for HF and closed-shell DFT methods

See Table 1 for a complete list of TeraChem capabilities.

---

<sup>1</sup> W. L. Jorgensen, J. Chandrasekha, J. D. Madura, R. W. Impey and M. L. Klein, *J. Chem. Phys.* **79** 926 (1983).

## Obtaining TeraChem

To purchase a copy of TeraChem visit <http://shop.petachem.com>. Pricing information is available at <http://www.petachem.com/pricing.html>.

## Citing TeraChem

Any published work that utilizes TeraChem shall include the following reference:

I.S. Ufimtsev and T.J. Martinez, Quantum Chemistry on Graphical Processing Units. 3. Analytical Energy Gradients and First Principles Molecular Dynamics, *J. Chem. Theory Comput.*, 2009, **5**, p2619.

Work which uses the geometry optimization or transition state finding utilities shall include the following reference:

J. Kästner, J.M. Carr, T.W. Keal, W. Thiel, A. Wander and P. Sherwood, DL-FIND: An Open-Source Geometry Optimizer for Atomistic Simulations, *J. Phys. Chem. A*, 2009, **113**, p11856.

Whenever dispersion corrections are used, the following references should be cited:

S. Grimme, J. Antony, S. Ehrlich, and H. Krieg, *J. Chem. Phys.*, 2010, **132**, p154104.

S. Grimme, S. Ehrlich, and L. Goerigk, *J. Comput. Chem.*, 2011, **32**, p1456.

## Acknowledgements

This software was developed by Ivan Ufimtsev and Todd Martinez at the University of Illinois at Urbana-Champaign and PetaChem, LLC. The authors would like to especially thank Nathan Luehr for contribution of the GPU accelerated code for construction of numerical DFT grids and Alexey Titov for contribution of GPU accelerated code for electron repulsion integrals involving d functions. Geometry optimization and transition state search calculations use the DL-FIND library created by Johannes Kästner at Stuttgart University. Hartree-Fock and DFT dispersion correction code is used with permission from Stefan Grimme at the University of Münster. TeraChem uses standard Gaussian basis sets available at EMSL website <http://www.emsl.pnl.gov/forms/basisform.html>

## II Getting started

### System requirements

This version of TeraChem was compiled and tested under 64-bit RedHat Enterprise Linux 5.5 operating system running on Intel Core2 quad-core and Intel Xeon 5520 dual quad-core CPU machines. An Nvidia compute capability 2.0 (Tesla C2050 or similar) or higher (i.e. Tesla K20 or similar) graphics card is required to run the program. Please refer to the CUDA Programming Guide at [http://www.nvidia.com/object/cuda\\_develop.html](http://www.nvidia.com/object/cuda_develop.html) for the most current list of Nvidia GPU's that meet this requirement. A CUDA driver (304.54 or later) must be installed on the system as well as v5.0 of the CUDA Toolkit. *Note that other CUDA toolkits, even newer versions, are not guaranteed to run correctly and will likely fail.* Details on how to obtain and install the CUDA driver are provided below.

Because the binary file is linked against the Intel MKL library, it is recommended to run TeraChem on Intel-based workstations.

The amount of CPU RAM needed depends on the size of the molecules that will be studied. If the molecules of interest are relatively small (less than 500 atoms), the usual 8Gb or 16Gb configuration is acceptable. For very large molecules (in excess of 10,000 basis functions), CPU RAM will often be a limiting factor. For example, molecules with 25,000 basis functions require almost 70GB of CPU memory.

### Installation

First, obtain and install the latest CUDA driver (TeraChem 1.5K requires 304.54 or later) and the CUDA 5.0 toolkit (choose the one appropriate for your Linux OS, e.g. RedHat Enterprise Linux 5.5 for RHEL 5.5 or CentOS 5.5) available for download free of charge at

```
https://developer.nvidia.com/cuda-downloads
```

If CUDA 5.0 is not installed on your machine and you do not have root permissions, you can install the toolkit in your home directory and set LD\_LIBRARY\_PATH environment variable accordingly:

```
$ export LD_LIBRARY_PATH=~/cuda/lib64:$LD_LIBRARY_PATH
```

Installation of the CUDA driver must be performed by a user with “root” permission, i.e. the superuser. Check with your local system administrator first, since adequate CUDA drivers may already be installed. Make sure you select the 64-bit Linux operating system. You will need to install the CUDA driver and the CUDA toolkit (ver. 5.0). The CUDA SDK is not required (but may be installed for other purposes if desired). After downloading the driver package, shut down the X server by typing

```
$ sudo init 3
```

Then launch the driver binary, and follow the instructions. Unpack the tc.tar archive using the following command (in a temporary directory which you may later remove):

```
$ tar xvf tc.tar
```

Run the install script by typing

```
$ chmod u+x install  
$ ./install
```

This script will verify that your machine has a suitable graphics card, verify that you accept the license terms, and install the software in a location of your choosing (the “TeraChem installation directory” or `instdir` in the following). The script will ask you for the location of the CUDA libraries (installed when the CUDA toolkit is installed, see above). It will also create a script that sets the appropriate environment variables. The `install` script will also put a temporary license file in place so you can begin using TeraChem immediately. However, this temporary license file is time-limited, so you will want to obtain a permanent license file. The `install` script ends with a form suitable for emailing to [help@petachem.com](mailto:help@petachem.com):

-----BEGIN HERE-----

Institution: \_\_\_\_\_

Ordered By: \_\_\_\_\_

MAC: 003048DB1D7E

IP: 171.64.125.189

-----END HERE-----

You can regenerate this at any time by typing

```
$ ./machid
```

in the TeraChem installation directory. Fill in the “Institution” and Ordered By” fields and email to [help@petachem.com](mailto:help@petachem.com). When we receive this from you, we will send you a permanent license file (which should be saved as `license.dat` in the TeraChem installation directory). TeraChem environment variable indicates where the licensing file can be found, i.e. `$TeraChem/license.dat`.

## Running sample jobs

TeraChem package contains several sample jobs located at

```
instdir/TeraChem/terachem/tests
```

After installation, `cd` to the `caffeine` directory and run TeraChem by typing

```
$ source instdir/TeraChem/SetTCVars.sh  
$ instdir/TeraChem/terachem start.sp
```

where `instdir` is the installation directory you chose during the install (defaults to your home directory) and `start.sp` is the name of a TeraChem input file. Note that the environment variable `TeraChem` is set by “source’ing” `SetTCVars.sh`. This is needed in order for TeraChem to locate the license and basis set library files.

The `start.sp` contains the required parameters of the job (including the filename of the file which contains the atomic coordinates for the molecule of interest). Most of the parameters have default values. The complete list of parameters available in this version is presented in Table 1. An example of the configuration file used for single point energy calculations of caffeine with the BLYP functional, DFT-D dispersion corrections, and the 6-31G\* basis set is:

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```
# Job: Single point energy of caffeine
#
# basis set
basis      6-31g*
# coordinates file
coordinates caffeine.xyz
# molecule charge
charge      0
# SCF method (rhf/b3lyp/etc...): DFT-BLYP
method      b3lyp
# add dispersion correction (DFT-D)
dfdt       yes
# type of the job (energy/gradient/md/minimize/ts): energy
run        energy
end
```

All lines beginning with the '#' character are considered comments and are ignored by TeraChem. There is no requirement on the line ordering in the start file except that the last line should be 'end'.

Below is the output from this example job. The program first lists the parameters followed by all GPUs used in the job. Each GPU has its amount of memory and compute capability printed next to it. The program then attempts to predict the maximum size of the molecule that can be handled on the current machine based on the amount of available CPU and GPU memory. *Note that the recommended maximum size of the system is approximate*. The SCF procedure, which includes the DIIS error (the maximum component of the DIIS error vector), integrated number of electrons, exchange-correlation energy, SCF energy, and the total time elapsed per iteration, completes the program's output.

```
$ ./terachem start

*****
*          TeraChem v1.5K
*          Hg Version: b133f5bd52e3+
*          Compiled for GeForce Cards
*          Chemistry at the Speed of Graphics!
*****
* This program may only be used in connection with
* a valid license from PetaChem, LLC. Use of this program
* or results therefrom indicates acceptance of all terms
* and conditions stated in the license and that a valid
* license agreement between the user and PetaChem, LLC
* exists. PetaChem, LLC does not warrant the correctness
* of results or their suitability for any purpose.
* Please email bugs, suggestions, and comments to
*          help@petachem.com
*
*****
*****
```

```
*****
* Compiled by toddmtz      Fri Dec 7 00:14:47 PST 2012 *
* Supported architectures: SM 1.3, 2.0, 3.0, 3.5
* Cuda compilation tools, release 5.0, V0.2.1221
*****
*****
```

```
Job started   Fri Dec 7 15:00:03 2012
On MTZ08 (available memory: 68820 MB)

##### RUNTIME INFO #####
./terachem start

NVRM version: NVIDIA UNIX x86_64 Kernel Module  304.54  Sat Sep 29 00:05:49 PDT 2012
GCC version:  gcc version 4.3.2 20081105 (Red Hat 4.3.2-7) (GCC)
```

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```
linux-vdso.so.1 => (0x00007ffffcddaf000)
libpthread.so.0 => /lib64/libpthread.so.0 (0x000000382e400000)
libcublas.so.5.0 => /usr/local/cuda-5.0/lib64/libcublas.so.5.0 (0x00007fe60e65c000)
libcufft.so.5.0 => /usr/local/cuda-5.0/lib64/libcufft.so.5.0 (0x00007fe60c656000)
libcuda.so.1 => /usr/lib64/libcuda.so.1 (0x00007fe60ba6f000)
libcudart.so.5.0 => /usr/local/cuda-5.0/lib64/libcudart.so.5.0 (0x00007fe60b815000)
libm.so.6 => /lib64/libm.so.6 (0x000000382dc00000)
libstdc++.so.6 => /usr/lib64/libstdc++.so.6 (0x000000383ce00000)
libgcc_s.so.1 => /lib64/libgcc_s.so.1 (0x0000003839600000)
libc.so.6 => /lib64/libc.so.6 (0x000000382d800000)
/lib64/ld-linux-x86-64.so.2 (0x000000382c400000)
libdl.so.2 => /lib64/libdl.so.2 (0x000000382e000000)
librt.so.1 => /lib64/librt.so.1 (0x000000382f400000)
libz.so.1 => /lib64/libz.so.1 (0x00000034c0a00000)
#####
***** License Check *****
* Use license from: /home/ufimtsev/petachem/TeraChem/license.dat
* Available Host id: 003048DA94D0
* Available Host id: 003048DA94D1
* Available Host id: 3243CCBF8B03
* License expires: 2013-01-05
***** License OK *****
Scratch directory: ./scr

XYZ coordinates caffeine.xyz
Using basis set: 6-31gs
Spin multiplicity: 1
Using DIIS algorithm to converge WF
WF convergence threshold: 3.00e-05
Maximum number of SCF iterations: 100
Incremental Fock matrix formation
Will switch to conventional Fock if diffuse functions are detected
PRECISION: DYNAMIC
X-matrix tolerance: 1.00e-04
DFT Functional requested: blyp
Method: BLYP with dispersion correction
Becke 1988 exchange functional: 1.0
Lee-Yang-Parr correlation functional: 1.0
Wavefunction: RESTRICTED

-----
|      DFTD3 V2.0 Rev 1      |
| S.Grimme, University Muenster   |
| Fri Dec 17 17:08:12 CET 2010    |
|          (Hawaii Version)     |
-----

Please cite DFT-D3 work done with this code as:
S. Grimme, J. Antony, S. Ehrlich and H. Krieg,
J. Chem. Phys., 132 (2010), 154104.
If used with BJ-damping cite also
S. Grimme, S. Ehrlich and L. Goerigk,
JCC, submitted.
For DFT-D2 the reference is
S. Grimme, J. Comput. Chem., 27 (2006), 1787-1799

DFT grid type: 1
Using dynamic DFT grids.
Initial guess generated by maximum overlap

*****
**** SINGLE POINT ENERGY CALCULATIONS ****
*****

using 2 out of 4 CUDA devices
Device 0:      Tesla K20c, 5119MB, CC 3.5 -- IDLE
Device 1:      Tesla C2050, 3071MB, CC 2.0 -- IDLE
Device 2:      GeForce GTX 690, 2047MB, CC 3.0 -- CPU THREAD 0
Device 3:      GeForce GTX 690, 2047MB, CC 3.0 -- CPU THREAD 1
-----
CPU Memory Available: 9008.38 MegaWords
GPU Memory Available: 255.91 MegaWords
Maximum recommended basis set size: 11200 basis functions
(limited by GPU memory)
-----
Using d-functions. Configuring GPUs accordingly.
0: CUBLAS initialized, available GPU memory: 1347MB
1: CUBLAS initialized, available GPU memory: 1346MB
```

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```
Basis set:      6-31gs
Total atoms:    24
Total charge:   0
Total electrons: 102 (51-alpha, 51-beta)
Number electrons modeled by ECPs: 0
Total orbitals: 230
Total AO shells: 104 (62 S-shells; 28 P-shells; 14 D-shells; 0 F-shells; 0 G-shells)
Spin multiplicity: 1
Nuclear repulsion energy (QM atoms): 933.356431723822 a.u.
```

```
Setting up the DFT grid...
time to set the grid = 0.01 s
DFT grid points: 23506 (979 points/atom)
Setting up the DFT grid...
time to set the grid = 0.03 s
DFT grid points: 65912 (2746 points/atom)
```

\*\*\* Start SCF Iterations \*\*\*

Iter	DIIS Error	Energy change	Electrons	XC Energy	Energy	Time(s)
>>> Purifying P... IDMP = 1.53e-14 <<<						
THRESPDP	set to 1.00e+00					
1	0.3143222065	-677.4535764850	101.9995043389	-89.9757230998	-677.4535764850	0.66
	>>> SWITCHING TO GRID 1 <<<					
2	0.6986364323	+6.5996837687	101.9970678470	-93.6280083063	-670.8538927162	0.41
3	0.2864124356	-8.2488710461	101.9980922169	-93.2210986394	-679.1027637623	0.39
4	0.1098582343	-0.9402443640	101.9979502441	-91.5162309843	-680.0430081263	0.41
5	0.1379180296	-0.0258751155	101.9976743328	-92.0184706858	-680.0688832418	0.40
6	0.0299984056	-0.1205332226	101.9979006529	-92.0222388541	-680.1894164644	0.40
7	0.0073251834	-0.0131962993	101.9978758774	-91.9713249086	-680.2026127637	0.39
	>>> SWITCHING TO GRID 1 <<<					
8	0.0036533335	-0.0012610982	101.9996482881	-91.9600999713	-680.2038738619	0.67
9	0.0022302215	-0.00001021822	101.9996468643	-91.9674933789	-680.2039760440	0.76
10	0.0005471869	-0.0000498203	101.9996473056	-91.9698688887	-680.2040258644	0.56
THRESPDP	set to 3.56e-03					
11	0.0002710918	+0.0000079054	101.9996488341	-91.9703058755	-680.2040179590	0.74
12	0.0001137490	-0.0000014169	101.9996487467	-91.9698501918	-680.2040193759	0.65
13	0.0000577038	-0.0000001696	101.9996487669	-91.9696944751	-680.2040195455	0.68
14	0.0000185995	-0.0000000001	101.9996487389	-91.9698173292	-680.2040195456	0.64

DISPERSION CONTRIBUTION TO ENERGY: -0.0562114195 a.u.

FINAL ENERGY: -680.2040195456 a.u.

CENTER OF MASS: {0.008017, 0.006143, 0.000066} ANGS

DIPOLE MOMENT: {3.430559, -0.664809, 0.000035} (|D| = 3.494382) DEBYE

Writing out molden info

Running Mulliken population analysis...

Total processing time: 7.94 sec

Job finished: Fri Dec 7 15:00:15 2012

## III TeraChem I/O file formats

### Input files

In addition to the start file containing job parameters, TeraChem requires two other files to start a job: coordinates and basis set.

### Atomic coordinates

The coordinates file fed to the coordinates parameter should be either in XMol or PDB format. In an XMol coordinates file, the first line specifies the number of atoms, and the second line provides a description of the system (it can be left blank). Atomic coordinates are listed starting from the third line. All coordinates are either in Angstroms (default) or Bohrs (this can be specified in the start file). Here is an example coordinates file for a hydrogen molecule:

```

2
Hydrogen Molecule - Xmol format
H 0.0      0.0      0.0
H 0.7      0.0      0.0

```

Some jobs (for example, transition state search using NEB method) require several sets of coordinates (frames). In this case all frames should be listed in the coordinates file one by one, i.e.

```

2
Hydrogen Molecule - Xmol format frame 1
H 0.0      0.0      0.0
H 0.7      0.0      0.0
2
Hydrogen Molecule - Xmol format frame 2
H 0.0      0.0      0.0
H 0.8      0.0      0.0

```

Note that there should be no blank lines between individual frames.

The PDB format often used for protein molecules is also supported and will be automatically assumed if the filename for the coordinates ends in ‘.pdb’. More details on PDB file format are available at <http://www.wwpdb.org/docs.html>

## Basis set file format

TeraChem supports standard atom-centered Gaussian type basis sets such as those available at EMSL website <http://www.emsl.pnl.gov/forms/basisform.html>. The basis set information used by TeraChem is provided by a set of files in `basis` directory. This directory should be located as `$TeraChem/basis`, where the environment variable `TeraChem` is set to the TeraChem installation directory by default. All basis sets are listed in individual files.

For every atom in a basis set, the first four lines specify the atom type, number of core orbitals, number of valence orbitals, and the total number of contracted basis functions centered on the atom. Consider an iron atom in STO-3G basis set.

```

ATOM Fe
COR 9
VAL 4
BFS 19

```

Here, `Fe` is an alias of the atomic nuclear charge, `z`, and `BFS` is the total number of atomic shells multiplied by the number of basis functions per shell (1 for s-, 3 for p-, 6 for d-shell, etc). The next block contains coefficients of atomic orbitals (AOs) that need to be precalculated separately for each atom and each basis set. Below are 2 out of 13 AOs of iron in STO-3G basis set.

```

 9.92667130e-01   1.93498346e-02   -1.55903797e-02   -7.99232061e-04   -0.00000000e+00
 -0.00000000e+00   -0.00000000e+00   -0.00000000e+00   -0.00000000e+00   -0.00000000e+00
 -0.00000000e+00   -0.00000000e+00   -0.00000000e+00   -0.00000000e+00   -0.00000000e+00
 -0.00000000e+00   5.90570755e-03   5.90570755e-03   5.90570755e-03
 ..... .
 0.00000000e+00   0.00000000e+00   0.00000000e+00   -2.44836578e-11   0.00000000e+00
 0.00000000e+00   0.00000000e+00   0.00000000e+00   0.00000000e+00   0.00000000e+00
 0.00000000e+00   0.00000000e+00   0.00000000e+00   -5.06687307e-01   -2.83366410e-01
 4.39883475e-01   -1.02744686e-02   -5.88180817e-01   5.98455286e-01

```

This information is required to construct the initial guess as described in

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J.-M. Langlois, T. Yamasaki, R.P. Muller, and W.A. Goddard III, Rule-Based Wave Function for Generalized Valence Bond Theory, *J. Phys. Chem.*, 1994, **98**, p13498.

In the AO block, core orbitals (COR) are listed first, followed by valence orbitals (VAL). Each atomic orbital thus is represented by BFS coefficients. Therefore, the total number of AO coefficients is equal to  $BFS^*(COR+VAL)$ . Basis shells, including the shells' type, number of primitive functions and corresponding contraction coefficients and exponents are listed at the end of the atom description.

```
S 3
 1447.4004110      0.1543289673
 263.6457916      0.5353281423
 71.35284019      0.4446345422
  * * * * *
P 3
 111.9194891      0.1559162750
 26.00768236      0.6076837186
 8.458505490      0.3919573931
  * * * * *
D 3
 6.411803475      0.2197679508
 1.955804428      0.6555473627
 0.754610151      0.2865732590
```

It should be noted that the ordering of basis functions in the AO expansion used to construct the initial guess must be the same as in the GTO block above. Within P-type shells, the basis functions are ordered as X, Y, Z, and within D-type shells as XY, XZ, YZ, XX, YY, ZZ.

It is straightforward to mix different basis sets for the same molecule. To do that, one needs to create a separate basis file which will contain all required basis functions. Here is an example calculation of a hydrogen molecule with STO-3G and 3-21G basis set used for the first and the second hydrogen atom, respectively:

```
ATOM  H1
COR 0
VAL 1
BFS 1
 1.00000000E+00

S 3
 3.4252509      0.154328967295
 0.6239137      0.535328142282
 0.1688554      0.444634542185

ATOM  H2
COR 0
VAL 1
BFS 2
 3.73407177E-01 7.17324343E-01

S 2
 5.4471780      0.156284978695
```

```

0.8245472    0.904690876670
S 1
0.1831916    1.000000000000

```

When mixing basis sets, the atom names in the coordinates file should be consistent with those provided in the basis files. Here is an example of the corresponding XYZ file.

```

2
Hydrogen Molecule
H1 0.0      0.0      0.0
H2 0.7      0.0      0.0

```

Note that a user can use a local copy of the basis directory if desired by setting the TeraChem environment variable appropriately. In that case, one should also ensure that a copy of the license.dat file exists in \$TeraChem.

## Fixing Atoms in Molecular Dynamics

It is possible to fix a number of atoms in molecular dynamics by listing the atom numbers in a file called `fixed_atoms`, located in the directory containing the input file. The format is simply the number of fixed atoms followed by the atom number (counting starts at zero) of each fixed atom, with one atom per line. The force on fixed atoms is simply zeroed out, with no modification to the forces on other atoms. Only QM atoms can be fixed in QM/MM simulations. This does not have any effect on optimizations.

## Ab Initio Steered Molecular Dynamics (AISMD)

Specific atoms can be subject to constant pulling forces to generate force-modified potential energy surfaces (or molecular dynamics on these FMPESS).<sup>2</sup> In order to accomplish this, one needs to specify the number of atoms subject to steering forces, the identity of the atoms which are being steered, the fixed (in Cartesian space) points they are being pulled toward, and the magnitude of the force. This is accomplished with the `steering` file (located in the same directory as the input file) which has the following format:

```

Number_Steered_Atoms
Steered_Atom_Index Fixed_Point_X Fixed_Point_Y Fixed_Point_Z Steering_Force

```

where the last line occurs Number\_Steered\_Atoms times. Coordinates of the fixed points should be given in bohr and the steering force should be given in atomic units.

## Output files

In addition to the information displayed on the screen, TeraChem creates several output files.

All calculations:

<sup>2</sup> M. T. Ong, J. Leiding, H. Tao, A. M. Virshup and T. J. Martínez, J. Amer. Chem. Soc. **131** 6377 (2009).

`scr/c0` (`scr/ca` and `scr/cb` in UHF and UKS jobs) – the converged WF binary file containing the MO coefficients `C[i][j]` where `i` (row) is the MO and `j` (column) is the AO basis function index. This file can be used as initial WF guess in subsequent calculations.

`scr/prjct` (`scr/prjcta` and `scr/prjctb` in UHF and UKS jobs) – the converged WF projected onto another (usually, larger) basis set. These files are generated by `project` jobs and used as an efficient initial guess.

`scr/orbitals.log` – the canonical MO orbitals in GAMESS format. This file name can be modified by the `orbitals` parameter. Because the orbitals require much disk space, they can be written every  $n^{\text{th}}$  MD step, specified by `orbitalswrtfrq`. The orbitals can be visualized by VMD. Note that the only meaningful information contained in this file is the MO coefficients. Everything else is printed for VMD to parse the file correctly.

`scr/molden.molf` – the canonical MO orbitals in Molden format.

`scr/charge.xls` – a tab-separated file containing Mulliken atomic charges.

`scr/oe.xls` – MO energies.

#### Geometry Optimization/Transition State Search:

`scr/optlog.xls` – a tab-separated file containing 7 columns of which only one (the first one) is currently used. The first column contains the SCF energy during geometry optimization or transition state search.

`scr/optim.xyz` – geometry optimization or transition state search (depending on the job type) trajectory file in the XMol format. The trajectory can be visualized by VMD.

#### Transition State Search:

`scr/neb_n.xyz` – an XMol file containing trajectory of the  $n^{\text{th}}$  NEB image. The last image (i.e. the `neb_10.xyz` if `min_image` equals 10) is the actual transition state that is also stored in `optim.xyz` file.

`scr/nebinfo` – contains energies of all (`min_image`-1) NEB images along the converged NEB path.

`scr/nebpath.xyz` – contains XYZ coordinates of all (`min_image`-1) NEB images along the converged NEB path.

#### MD simulation:

`scr/log.xls` – a tab separated file containing 7 columns: 1) SCF energy, 2) currently not in use, 3) Kinetic energy, 4) Temperature, 5) Total energy (SCF + Kinetic), 6) HOMO energy, 7) LUMO energy. All energies are in Hartree and the temperature is in degrees Kelvin. In NVT dynamics, the Total energy does not include the contribution from the damping force, and thus should not be conserved.

`scr/coors.xyz` – the MD trajectory geometry file in the XMol format. The trajectory can be visualized by VMD.

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`scr/vel.log` – contains atomic velocities along in the MD trajectory. The format is similar to that of `scr/coors.xyz` except there is a blank line separating each set of velocities.

`scr/restart.md` – a binary MD restart file containing all information (wavefunction, coordinates, velocities, etc) required to restart an MD job. By default, this information is stored at every 100<sup>th</sup> MD iteration. A user can change it by specifying `restartmdfreq` parameter in the start file.

## IV More details on some TeraChem capabilities

### Wavefunction projection

Sometimes, especially when transition metals or diffuse basis functions are present in a system, the SCF procedure does not converge due to insufficiently accurate initial guess. In such cases, it often helps to converge the wavefunction using a smaller basis set (mini, sto-3g, etc) with no diffuse functions, then project the solution onto the desired basis set and start another SCF procedure using the projected wavefunction as initial guess. Below is an example of such project job. Here, the calculations are performed in a smaller (sto-3g) basis set, and the converged wavefunction is projected onto 6-31g\*\* basis set.

```
run          project
basis        sto-3g
projectbasis 6-31g**
charge       0
coordinates   water.xyz
end
```

The projected wavefunction is stored in `scr/prjct` file.

### MD with spherical boundary conditions

In an MD simulation, it is possible to impose spherical boundary conditions to prevent “evaporation” events or constrain a system to a given density. The spherical boundary conditions are provided in the form of a sum of two harmonic terms,

$$U_{\text{constr}}(r) = k_1((r - R_{\text{center}}) - R_1)^2 + k_2((r - R_{\text{center}}) - R_2)^2,$$

where  $R_{\text{center}}$  is the center of mass of the system calculated for the first MD frame and then fixed. By default,  $k_1$  is set to 10.0 kcal/(mol Å<sup>2</sup>) and  $k_2$  is set to zero. The simplest way to impose spherical boundary conditions is to set

```
mdbc        spherical
md_density  1.0
```

in the start file. `md_density` specifies the density of the system in g/mL used to automatically adjust  $R_1$ . If `md_density` is not provided,  $R_1$  needs to be set explicitly using the `md_r1` keyword. Note that constraining forces are not applied to hydrogen atoms.

### QM/MM Functionality

Limited QM/MM functionality is available in this release. Only water molecules are available for MM treatment and they will be modeled using the TIP3P force field. One signals that a QM/MM calculation is desired by including the `qm/mm` keyword in `inputfile`. The keyword specifies the location of a file with coordinates for the desired MM water molecules. Note that the coordinates of the MM waters are *not* in the `coordinates` file. The water molecule positions must appear in a specific order – grouped by water molecule and in the order O,H,H. We provide an example for one QM water molecule surrounded by one MM water molecule.

`inputfile:`

```

basis      6-31g
qmmm      mmwater.xyz
coordinates qmwater.xyz
run        minimize
end

```

qmwater.xyz:

3

O	6.144353	-0.788526	-6.483525
H	6.378922	-1.564817	-5.989856
H	6.475062	-0.043630	-5.995619

mmwater.xyz:

3

O	6.450776	0.962914	-3.536427
H	6.338156	1.514355	-2.771897
H	6.519847	1.524502	-4.303300

It is also possible to use TeraChem with Amber 12 for more involved QM/MM calculations, e.g. where the MM region includes protein residues. See the Amber 12 manual<sup>3</sup> for details.

## Integration with NBO6

This release of TeraChem is interfaced with the NBO6 package of Weinhold and coworkers.<sup>4</sup> NBO deletions are not currently supported, nor are analyses which require the matrix elements of the dipole operator. However, many of the usual natural bond orbital (NBO) and natural population analyses (NPA) are supported. Setting the `nbo` keyword in the `startfile` to `npa` or `full` will give natural population analysis or full NBO/NPA analysis. You may also use advanced keywords (documented in the NBO manual) if desired. In this case, the `nbo` keyword should be set to `advanced` and the `startfile` should contain a `$nbo` group as documented in the NBO manual. An example of this usage of advanced keywords for CH<sub>3</sub>NH<sub>2</sub> is provided in `TeraChem/tests/nbo`. Note that the `NBOEXE` environment variable needs to point to the NBO6 executable file in order for successful integration with TeraChem. This is currently set in the `SetTCVars.sh` script. The NBO6 executable is included with the TeraChem distribution and resides in the TeraChem directory.

## Ghost Atoms

Sometimes it is desirable to do calculations including the basis functions of an atom (or atoms) without including the nucleus or electrons corresponding to the atom. This is most often used to

<sup>3</sup> <http://ambermd.org>

<sup>4</sup> <http://www.chem.wisc.edu/~nbo5>

calculate basis set superposition error (BSSE). For example, the Boys-Bernardi counterpoise correction<sup>5</sup> for the interaction energy of fragments A and B is:

$$E_{AB}^{counterpoise} = E_{AB}(H_{AB}, R_A, R_B; A \cup B) - E_A(H_A, R_A; A \cup B) - E_B(H_B, R_B; A \cup B)$$

where the first arguments in the parentheses are the Hamiltonian and coordinates it depends on and the second arguments (after the semicolon) are the basis sets used. Thus, in the second term on the right, the calculation has the atoms of A (and the corresponding basis functions), but also includes the basis functions for fragment B (but *not* the electrons or nuclei corresponding to the atoms of B). In TeraChem, one can include ghost atoms (atoms which contribute basis functions to the calculation, but *not* electrons or nuclear charges) by prefacing the element name with the letter X. For example, the following is a coordinates (XYZ) file used in calculating the counterpoise-corrected interaction energy for water dimer:

6

O	-0.029481	-0.086991	-0.533710
H	-0.660044	-0.772698	-0.712077
H	0.086741	0.049151	0.406478
XO	0.306708	0.307377	2.262155
XH	1.109626	0.019913	2.679671
XH	-0.049150	1.088298	2.668550

The last three atoms will be “ghost” or “dummy” atoms, i.e. the system has only 10 electrons, one oxygen atom nucleus and two hydrogen atom nuclei, but basis functions will be used on all six listed atoms (according to the basis set listed in the start file).

## Geometry Optimization and Transition State Search

The `instdir/tests/sp` directory contains a simple configuration file (`start.go`) used for geometry optimization of a spiropyran molecule:

```
# basis set
basis      6-31g
# coordinates file
coordinates sp.xyz
# molecule charge
charge      0
# SCF method (rhf/blyp/b3lyp/etc...): RHF
method      rhf
# type of the job (energy/gradient/md/minimize/ts): geometry
# optimization
run        minimize
end
```

The optimization is triggered by the ‘minimize’ keyword. Note that the `sp.xyz` file in fact contains two sets of coordinates (frames) one by one. In geometry optimization jobs, only the first frame is taken into account while the others (if any) are ignored. The second frame,

---

<sup>5</sup> S. F. Boys and F. Bernardi, Mol. Phys. **19** 552-566 (1970)

however, is required by transition state search jobs and represents the second NEB endpoint in NEB calculations. The TS search is triggered by the ‘ts’ keyword, i.e. file (start.ts)

```
# basis set
basis      6-31g
# coordinates file
coordinates sp.xyz
# molecule charge
charge      0
# SCF method (rhf/b3lyp/b3lyp/etc...): RHF
method      rhf
# type of the job (energy/gradient/md/minimize/ts): TS search
run        ts
end
```

\$constraints ... \$end group in the configuration file allows one to impose geometrical constraints during optimization (frozen atoms, fixed bond lengths, angles, and dihedrals). instdir/tests/constraints directory contains two example start files for constrained geometry optimization jobs.

Constrained geometry optimization can be performed in any type of coordinates with two exceptions: 1) if DLC coordinates are used for optimization (the default), no atoms may be frozen and 2) if Cartesian coordinates are used for optimization (`min_coordinates cartesian`), only frozen atom constraints are allowed. Below is an example of a job configuration start file. Note that all constraints should be listed in separate lines, and enumeration of atoms begins from 1. Constrained coordinates will be constrained to the value they have in the starting geometry.

```
# basis set
basis      6-31g
# coordinates file
coordinates C10H22.inp
# molecular charge
charge      0
# optimize geometry
run        minimize
end

$constraints
# bond connecting atoms 32 and 29
bond      32      29
# 32-29-26 angle
angle      32      29      26
# 32-29-26-27 dihedral
dihedral   32      29      26      27
$end
```

Finally, it is straightforward to freeze all hydrogen or non-hydrogen atoms by `atom hydrogens` or `atom heavy`, respectively.

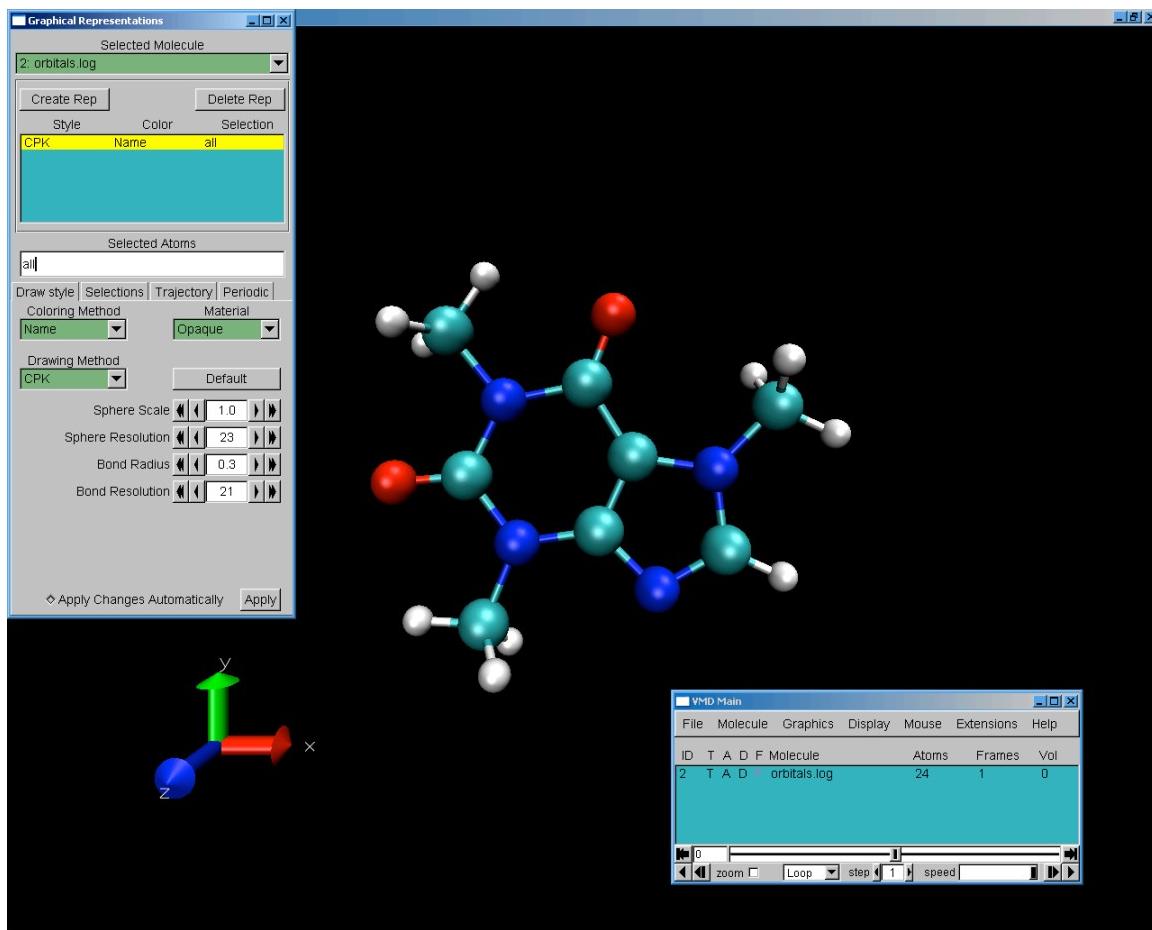
## V Trajectory visualization

To create a simple MD trajectory, you can use the example in `instdir/tests/caffeine`. The configuration file `start.md` is:

```
# basis set
basis          6-31g
# coordinates file
coordinates    caffeine.xyz
# molecule charge
charge         0
# SCF method (rhf/blyp/b3lyp/etc...): DFT-BLYP
method         blyp
# type of the job (energy/gradient/md/minimize/ts): MD
run           md
# number of MD steps
nstep         10
# dump orbitals every MD step
orbitalswrtfrq 1
end
```

The only difference between this file and the file used for the single point calculations are the `nstep` and `orbitalswrtfrq` parameters, which are now set to 10 and 1, respectively. The keyword `orbitalswrtfrq` ensures that the molecular orbitals will be written at every MD step (out of 10, total).

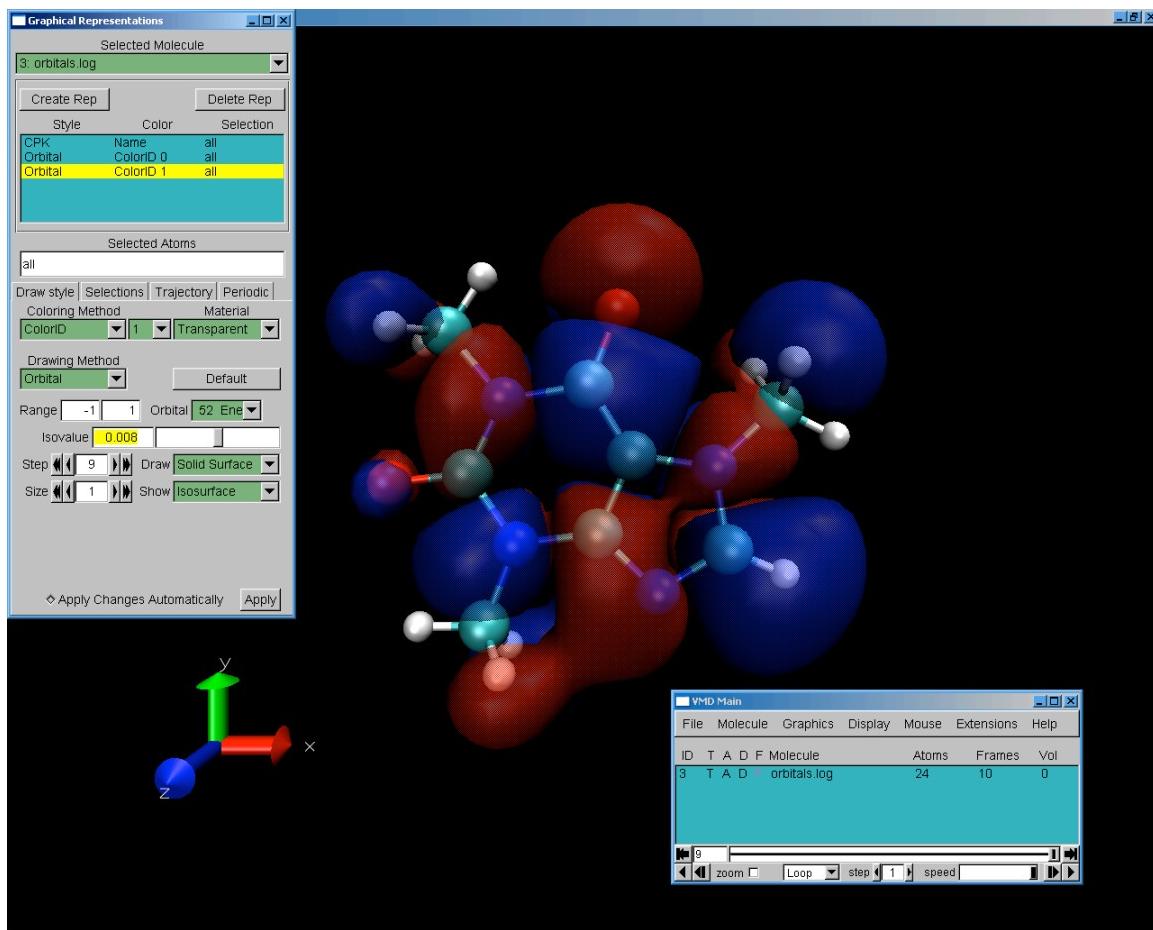
The TeraChem output coordinates file format is compatible with VMD. To open the trajectory file, go to **VMD** → **File** → **New Molecule**. Browse to the output `coors.xyz` file and make sure that the file type is correctly determined. Otherwise, select **XYZ** in the [**Determine file type**] menu. After the trajectory is loaded, you can adjust the representation settings so that the final molecule looks as shown in Figure 1.



**Figure 1.** Caffeine molecule geometry output (`coors.xyz`) in VMD.

The molecular orbitals can be visualized with VMD in similar fashion. Again, go to VMD → File → New Molecule. Browse to the output `orbitals.log` file and make sure that the determined file type is GAMESS. Otherwise, select GAMESS in the [Determine file type] menu. To display both positive and negative isosurfaces you will need to create two representations for each orbital (one for the positive isosurface and one for the negative isosurface). After adjusting the atomic representation, create a new one and select “orbital” in the [Drawing Method] menu. All MO’s will be listed in the [Orbital] dropdown along with the orbital energy. The [Isovalue] scroll bar controls the isosurface value (in a.u.).

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**Figure 2.** Caffeine molecule orbitals output (`orbitals.log`) in VMD.

## VI Interactive calculations

Interactive calculations are especially suitable for remote jobs when TeraChem is running on a remote machine (cluster) and the trajectory visualization is performed on a local desktop. TeraChem can visualize the geometry optimization, TS search, and molecular dynamics trajectories in real time, i.e. interactively as the calculations run. These interactive molecular dynamics (IMD) runs are practical for molecules with up to approximately 20 atoms using a hardware solution with eight GPUs. Future versions will also allow for user manipulation of the molecule, i.e. imposing external forces on atoms. In the `instdir/tests/benzene` directory, you will find the files needed to try IMD for benzene. The configuration file `start.imd` reads:

```
# basis set
basis      sto-3g
# coordinates file
coordinates C6H6.pdb
# molecule charge
charge      0
# SCF method (rhf/blyp/b3lyp/etc...): Restricted Hartree-Fock
method      rhf
# type of the job (energy/gradient/md/minimize/ts): MD
run        md
# initial temperature in K
tinit      1000
# this triggers interactive molecular dynamics
# imd specifies the port VMD should connect to
imd        54321
# number of MD steps
nstep      1000
end
```

Open two terminal windows. In the first window, launch VMD and load in the coordinates of the benzene molecule from `C6H6.pdb`. In the second terminal window, launch TeraChem, which will initialize the simulation and pause for connection to be made with VMD. Now, in VMD, select **Extensions**→**Simulation**→**IMD Connect (NAMD)**. Type `localhost` (or the IP address of the remote machine on which TeraChem is currently running) in the **Hostname** field and `54321` (the port specified by the `imd` keyword in the TeraChem input file) in the **Port** field. Click the **Connect** radio button and you should see TeraChem executing in its window while the benzene molecule vibrates in the VMD display window.

## VII TeraChem job parameters

Available job parameters are listed in Table 1. Note the use of the character “|” which should be interpreted as “or.” For example, `x | y` means that *one* of x or y should be entered.

**Table 1.** Available TeraChem job parameters.

Parameter	Description	Default value
General parameters		
<code>scrdir</code>	Scratch directory	<code>./scr</code>
<code>run</code>	Sets calculation type <code>energy</code> – Single point energy <code>project</code> – Wavefunction projection for better initial guess <code>gradient</code> – Energy and gradient <code>minimize</code> – Optimize geometry <code>ts</code> – Transition state search <code>md</code> – Born-Oppenheimer Molecular Dynamics	<code>energy</code>
<code>gpus</code>	Number of GPUs to use in parallel. Sometimes one needs to change the default order of the devices. For example when device zero is used solely for display purposes. In such cases, the following line specifies which GPUs should be used (the enumeration starts from 0):  <code>gpus 3 1 3 2</code> or <code>gpus 3</code> to use the default device order <code>0 1 2</code>	<code>all</code>
<code>precision</code>	<code>single   mixed   dynamic   double</code>  By default, TeraChem uses a dynamic precision scheme where two-electron integrals larger than some threshold value are computed with double precision and the rest are computed with single precision. The threshold which determines the splitting between single precision and double precision evaluation of the integrals is determined automatically during the SCF calculation. The mixed precision scheme uses a static threshold to determine which integrals should be evaluated in double precision (those larger than <code>threspdः</code> , which can be set in <code>inputfile</code> ). Users can	<code>dynamic</code>

	also request to use full single or double precision depending on the job purposes. Note that accumulation of all quantities on GPU is <i>always</i> performed with double precision accuracy to avoid floating point summation errors.	
threspdp	Threshold used to split single and double precision work in mixed precision calculations (atomic units).	0.001
coordinates	Name of file containing atomic coordinates	not set
qmmm	Name of file containing MM water coordinates  The presence of this keyword triggers a QM/MM calculation	not set
basis	All available basis sets are located in <code>basis/</code> directory. In addition, users can construct customized basis sets. Here are examples of correct entries:  <code>sto-3g, 6-31++G**, 6-311++G(3df,3pd),</code> <code>aug-cc-pvdz, mini(s), midi!</code>  When the string is parsed, all '*' symbols are replaced by 's' and all parenthesis are replaced by brackets to comply with Unix file naming rules.	not set
projectbasis	In <code>project</code> jobs, the converged wave function is projected onto <code>projectbasis</code> and stored in <code>scr/prjct</code> file ( <code>scr/prjcta</code> <code>scr/prjctb</code> in unrestricted methods).	not set
charge	The total charge of the molecule (integer).	not set
spinmult	Spin multiplicity, $2S+1$ , of wavefunction (integer).	1
method	Restricted, unrestricted, and restricted open shell HF and KS.  Restricted wavefunction: <code>rhf svwn b3lyp b3lyp1 b3lyp5 pbe revpbe pbe0 revpbe0 wpbe wpbeh bop mubop camB3LYP b97 wb97 wb97x</code>  <code>b3lyp</code> or <code>b3lyp1</code> : VWN1 correlation <code>b3lyp5</code> : VWN5 correlation Unrestricted calculations are invoked by adding 'u' prefix, i.e. <code>uhf ub3lyp ub3lyp, etc</code>  Restricted open shell calculations are invoked by adding 'ro' prefix, i.e. <code>rohf rob3lyp rob3lyp, etc</code>	rhf

<code>rc_w</code>	Range correction scaling parameter $\omega$ required by DFT functionals with a fraction of exact long range exchange operator	depends on DFT functional
<code>c_ex</code>	Long range exact exchange operator contribution	depends on DFT functional
<code>dftgrid</code>	Integer value within [0-5] range, inclusive. Larger numbers are denser grids (and hence provide more accurate results). Grid 0 contains ~800 grid points/atom and grid 5 contains ~80,000 points per atom. The default grid (type 1) contains about 3,000 points per atom.	1
<code>dynamicgrid</code>	<code>yes no</code>  This parameter enables use of dynamical DFT grids. When it is on, grid 0 is used to converge the wavefunction until the DIIS error reaches the <code>gridthre</code> value (default: 0.01). Grid <code>dftgrid</code> is then used to finally converge the wavefunction.	<code>yes</code>
<code>gridthre</code>	Threshold for switching dynamical DFT grids.	0.01
<code>guess</code>	<code>generate path/to/the/WFfile</code>  <code>generate</code> means the initial WF guess is generated from scratch using maximum orbital overlap; <sup>6</sup> otherwise, it is loaded from the WF file. The WF is dumped in the end of each calculation to the <code>scr/c0</code> file ( <code>scr/ca scr/cb</code> in unrestricted methods).  To load WF in unrestricted calculations, <code>guess</code> requires two parameters, for example  <code>guess scr/ca scr/cb</code>	<code>generate</code>
<code>scf</code>	<code>diis</code> – Use Pulay's DIIS <sup>7</sup> for SCF convergence  <code>oda</code> – Use optimal damping <sup>8</sup> for SCF convergence  <code>diis+a</code> – Hybrid DIIS/A-DIIS <sup>9</sup> scheme  <code>diis_prime</code> – DIIS with density matrix purification <sup>10</sup> instead of diagonalization (can be faster when there are more than 3000 basis functions)	<code>diis</code>

<sup>6</sup> J.-M. Langlois et al, J. Phys. Chem. **98**, 13498 (1994).<sup>7</sup> P. Pulay, J. Comp. Chem. **3**, 556 (1982).<sup>8</sup> E. Cancès and C. Le Bris, Int. J. Quantum Chem. **79**, 82 (2000).<sup>9</sup> X. Hu and W. Yang, J. Chem. Phys. **132**, 054109 (2010).

start_diis	When using optimal damping, switch to DIIS once DIIS error falls below threshold (float).	0.01
fock	incremental – Force incremental Fock matrix formation  conventional – Force conventional Fock matrix formation  auto – Use conventional algorithm if diffuse basis functions are detected or incremental otherwise	auto
maxit	Maximum number of SCF iterations (integer).	100
convthre	WF convergence threshold (float).	3.0e-5
threall	Two-electron integral threshold (float). Two electron integrals less than this threshold are neglected.	10 <sup>-11</sup>
xtol	Basis set linear dependency threshold (float). When diffuse basis functions are used, <code>xtol</code> and <code>convthre</code> may need to be raised up to ~1.0e-3	1.0e-4
dispersion dftd (alias)	Should empirical dispersion corrections be used?  yes no d3 d2  D2 <sup>11</sup> and D3 <sup>12</sup> are two different dispersion parameterizations developed by S. Grimme.	no
units	Units used for coordinates ( <code>angstrom</code> or <code>bohr</code> )	angstrom
nbo	NBO analysis (no yes npa full advanced \$nbo)  no – no NBO analysis  yes npa – Natural population analysis only  full – Full NPA/NBO analysis  advanced \$nbo – Analysis with \$NBO input  TeraChem uses v6.0 of the NBO package. See <a href="http://www.chem.wisc.edu/~nbo5">www.chem.wisc.edu/~nbo5</a> for documentation of the earlier v5.0.	no
polarizability	Calculate polarizability tensor? (yes no)	no
safemode	Allow to pre-configure GPUs for basis sets with d-functions. Setting this parameter to ‘no’ bypasses pre-configuration step but allows to have more GPU	yes

<sup>10</sup> A. M. N. Niklasson, Phys. Rev. B **66**, 155115 (2002).<sup>11</sup> S. Grimme, J. Comp. Chem. **27**, 1787 (2006).<sup>12</sup> S. Grimme, J. Antony, S. Ehrlich and H. Krieg, J. Chem. Phys. **132**, 154104 (2010)

	memory (e.g. GeForce). Use at your own risk.	
<b>Geometry Optimization and Transition State Search</b>		
min_print	How much output is desired (something verbose debug)	verbose
nstep	Maximum number of optimization/TS search steps	100
min_tolerance	Termination criterion based on the maximum energy gradient component	$4.5 \times 10^{-4}$
min_tolerance_e	Termination criterion based on the SCF energy change	$10^{-6}$
min_coordinates	Type of coordinates in which optimization/TS search is performed (cartesian dlc dlc_tc hdlc hdlc_tc)	dlc
min_method	Optimization/TS search method (sd cg1 cg2 lbfgs)  sd – steepest descent  cg1 and cg2 – conjugate gradient  lbfgs – L-BFGS	lbfgs
min_hess_update	Hessian update algorithm (never powell bofill bfsgs)  If never, the Hessian is recalculated using finite differences at each step	bfsgs
min_init_hess	Initial Hessian (fischer-almlof one-point two-point diagonal identity)  one-/two-point – exact Hessian from finite differences  diagonal – only diagonal elements are calculated using final differences, off-diagonal elements equal zero  identity – initial Hessian is an identity matrix  fischer-almlof – chemically-motivated guess Hessian, usually the best choice	fischer-almlof
min_delta	Atomic displacement in finite difference calculations	0.003
min_max_step	Maximum step size in internal coordinates	0.5
min_restart	Whether the optimization/TS search job is started from scratch (no) or loaded from the checkpoint files (yes)	no

min_dump	How often the checkpoint files are written. By default at each 10 <sup>th</sup> step.	10
ts_method	<p>Transition state search method            (neb_free neb_restricted neb_frozen             neb_free_cart neb_restricted_cart             neb_frozen_cart)</p> <p>neb_free – Nudged Elastic Band (NEB) with free endpoints</p> <p>neb_restricted – NEB with endpoints allowed to move perpendicularly to their tangent direction</p> <p>neb_frozen – NEB with frozen endpoints</p> <p>neb_x_cart – only initialization is performed in min_coordinates coordinates, while the TS search is done in Cartesians</p>	neb_free
min_image	<p>Number of NEB images in the TS search calculations. Should be greater than one. The images are listed in the input coordinates file (specified by coordinates). If the number of images found is smaller than min_image, the program will automatically generate missing images by interpolation. At least two images (endpoints) should be listed in the coordinates file. The last one (i.e. the min_image<sup>th</sup>) is the climbing image.</p>	10

### Molecular dynamics parameters

nstep	Total number of MD steps. Set nstep to 0 for single-point energy calculations (integer).	10 <sup>6</sup>
integrator	<p>The way the initial WF guess is generated in subsequent MD iterations. In the first several iterations the initial guess is constructed from scratch.            (reversible_d reversible regular reset)</p> <p>reversible_d: time-reversible integrator with dissipation<sup>13</sup></p> <p>reversible: time-reversible integrator without dissipation<sup>14</sup></p> <p>regular: initial WF guess is taken from converged</p>	reversible_d

<sup>13</sup> A.M.N. Niklasson et al, J. Chem. Phys. **130**, 214109 (2009).

<sup>14</sup> A.M.N. Niklasson et al, Phys. Rev. Lett. **97**, 123001 (2006).

	WF at previous MD step  reset: initial WF guess is generated from scratch at each MD step	
rseed	Seed for random number generator. Set this to a different integer for different MD runs (or set it to a known seed in order to reproduce an earlier run).	1351351
timestep	MD integration time step in femtoseconds (float)	1.0
thermostat	Temperature control – velocity rescaling or Langevin dynamics (rescale or langevin)	rescale
rescalefreq	When velocity rescaling is used, determines how often the velocities are rescaled. For instance, setting rescale to 1000 will force rescaling at every 1000 <sup>th</sup> MD step. To obtain NVE dynamics, set rescalefreq to a value larger than nstep.	$2 \cdot 10^9$
tinit	Initial temperature (K) sampled from Boltzmann distribution of velocities at T = tinit (float)	300.0
t0	Thermostat temperature (K) (float)	300.0
lnvtime	The Langevin damping time (fs), only used when thermostat is set to langevin.	1000.0
orbitals	Path to an output file containing the canonical molecular orbitals. The orbitals are printed in GAMESS format and can be visualized by VMD.	orbitals.log
molden	Path to an output file containing the canonical molecular orbitals in Molden format	molden.molf
orbitalswrtfrq	Determines how often the orbitals are written to the output file. Due to large size (sometimes the orbitals require 100MB and even more of disk space) it does not make sense to write orbitals at every MD iteration. (integer)	$2 \cdot 10^9$
velocities	Filename containing initial velocities (or random to specify that initial velocities should be drawn from Boltzmann distribution)	random
restartmd	Path to the MD restart binary file. If set, the data in this file (rather than coordinates, velocities, temperature, etc) is used to start MD trajectory.	not set
restartmdfreq	Specifies how often the restart data is dumped to scr/restart.md	100
mdbc	spherical  Enables spherical boundary conditions	not set

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md_density	If set, R <sub>1</sub> is automatically adjusted to the specified density in g/mL	1.0
md_r1, md_r2	R <sub>1</sub> and R <sub>2</sub> parameters for spherical boundary conditions in Å	0.0, 0.0
md_k1, md_k2	k <sub>1</sub> and k <sub>2</sub> parameters for spherical boundary conditions in kcal/(mol Å <sup>2</sup> )	10.0, 0.0
imd	Invokes interactive molecular dynamics by specifying the corresponding VMD port	not set

## **Contact information**

We will be pleased to receive any feedback on your experience with TeraChem. Should you have any suggestions, concerns, or bug reports, please email them to [help@petachem.com](mailto:help@petachem.com). Also, please feel free to participate in the TeraChem User Forum at <http://www.petachem.com/forum>. TeraChem developers monitor this forum frequently and are happy to answer questions about the software.

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## Release Notes

v1.50K:

- Fixed memory allocation problem which caused CUDA errors when running on multiple cards with shells that had few basis functions, e.g. when running with a single set of d functions over four cards.
- Implemented workaround for NVIDIA's texture bug (which caused Tesla C2075 and GeForce 580 GTX to hang randomly). Although we still do not recommend these cards, all tests so far suggest they can run TeraChem without errors.
- Document the ability to use ghost atoms (atoms that have basis functions but no electrons or nucleus – used in computing basis set superposition error).
- Document the ability to use matrix purification instead of diagonalization.
- Added A-DIIS/DIIS convergence scheme